

# Union Calendar No. 47

118TH CONGRESS  
1ST SESSION

# H. R. 467

**[Report No. 118-67, Part I]**

To amend the Controlled Substances Act with respect to the scheduling  
of fentanyl-related substances, and for other purposes.

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## IN THE HOUSE OF REPRESENTATIVES

JANUARY 24, 2023

Mr. GRIFFITH (for himself, Mr. LATTA, Mrs. RODGERS of Washington, Mr. GUTHRIE, Mr. BILIRAKIS, Mr. BUCSHON, Mr. HUDSON, Mr. BURGESS, Mr. CARTER of Georgia, Mr. DUNCAN, Mr. DUNN of Florida, Mr. CRENSHAW, Mr. JOYCE of Pennsylvania, Mr. BALDERSON, Mrs. HARSHBARGER, Mrs. MILLER-MEEKS, Mrs. CAMMACK, Mr. ALLEN, Mr. WALBERG, Mr. CURTIS, Mr. PALMER, Mr. BUCHANAN, Mr. BANKS, Mr. FITZGERALD, and Mr. MOONEY) introduced the following bill; which was referred to the Committee on Energy and Commerce, and in addition to the Committee on the Judiciary, for a period to be subsequently determined by the Speaker, in each case for consideration of such provisions as fall within the jurisdiction of the committee concerned

MAY 17, 2023

Additional sponsors: Mr. PFLUGER, Mr. ARMSTRONG, Mr. PENCE, Mr. AUSTIN SCOTT of Georgia, Mr. OBERNOLTE, Mr. WOMACK, Mr. EDWARDS, Mr. FINSTAD, Mr. CARL, Mrs. LESKO, Mr. MIKE GARCIA of California, Mr. CISCOMANI, Mr. CLINE, Mrs. BICE, Mr. CARTER of Texas, Mr. MOOLENAAR, Mr. ROGERS of Kentucky, Mr. BEAN of Florida, Mr. MEUSER, Mr. BOST, Mr. BARR, Mr. VAN DREW, Mr. D'ESPOSITO, Mrs. MILLER of West Virginia, Mrs. CHAVEZ-DEREMER, Mr. WESTERMAN, Mr. LANGWORTHY, and Mr. BURCHETT

MAY 17, 2023

Reported from the Committee on Energy and Commerce with an amendment

[Strike out all after the enacting clause and insert the part printed in italic]

MAY 17, 2023

Committee on the Judiciary discharged; committed to the Committee of the Whole House on the State of the Union and ordered to be printed

[For text of introduced bill, see copy of bill as introduced on January 24, 2023]

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## A BILL

To amend the Controlled Substances Act with respect to the scheduling of fentanyl-related substances, and for other purposes.

1       *Be it enacted by the Senate and House of Representa-*  
2   *tives of the United States of America in Congress assembled,*  
3   **SECTION 1. SHORT TITLE.**

4       *This Act may be cited as the “Halt All Lethal Traf-*  
5   *ficking of Fentanyl Act” or the “HALT Fentanyl Act”.*

6   **SEC. 2. CLASS SCHEDULING OF FENTANYL-RELATED SUB-**  
7                   **STANCES.**

8       *Section 202(c) of the Controlled Substances Act (21*  
9   *U.S.C. 812(c)) is amended by adding at the end of schedule*  
10   *I the following:*

11       “(e)(1) Unless specifically exempted or unless listed in  
12   another schedule, any material, compound, mixture, or  
13   preparation which contains any quantity of a fentanyl-re-  
14   lated substance, or which contains the salts, isomers, and  
15   salts of isomers of a fentanyl-related substance whenever the  
16   existence of such salts, isomers, and salts of isomers is pos-  
17   sible within the specific chemical designation.

18       “(2) For purposes of paragraph (1), except as provided  
19   in paragraph (3), the term ‘fentanyl-related substance’  
20   means any substance that is structurally related to fentanyl  
21   by 1 or more of the following modifications:

22       “(A) By replacement of the phenyl portion of the  
23   phenethyl group by any monocycle, whether or not  
24   further substituted in or on the monocycle.

1           “(B) By substitution in or on the phenethyl  
2       group with alkyl, alkenyl, alkoxy, hydroxyl, halo,  
3       haloalkyl, amino, or nitro groups.

4           “(C) By substitution in or on the piperidine  
5       ring with alkyl, alkenyl, alkoxy, ester, ether,  
6       hydroxyl, halo, haloalkyl, amino, or nitro groups.

7           “(D) By replacement of the aniline ring with  
8       any aromatic monocycle whether or not further sub-  
9       stituted in or on the aromatic monocycle.

10          “(E) By replacement of the N-propionyl group  
11       with another acyl group.

12          “(3) A substance that satisfies the definition of the  
13       term ‘fentanyl-related substance’ in paragraph (2) shall  
14       nonetheless not be treated as a fentanyl-related substance  
15       subject to this schedule if the substance—

16           “(A) is controlled by action of the Attorney Gen-  
17       eral under section 201; or

18           “(B) is otherwise expressly listed in a schedule  
19       other than this schedule.

20          “(4)(A) The Attorney General may by order publish  
21       in the Federal Register a list of substances that satisfy the  
22       definition of the term ‘fentanyl-related substance’ in para-  
23       graph (2).

24          “(B) The absence of a substance from a list published  
25       under subparagraph (A) does not negate the control status

1 *of the substance under this schedule if the substance satisfies*  
2 *the definition of the term ‘fentanyl-related substance’ in*  
3 *paragraph (2).”.*

4 **SEC. 3. REGISTRATION REQUIREMENTS RELATED TO RE-**

5 ***SEARCH.***

6 (a) *ALTERNATIVE REGISTRATION PROCESS FOR*  
7 *SCHEDULE I RESEARCH.—Section 303 of the Controlled*  
8 *Substances Act (21 U.S.C. 823) is amended—*

9 (1) *by redesignating the second subsection (l) (re-*  
10 *lating to required training for prescribers) as sub-*  
11 *section (m); and*

12 (2) *by adding at the end the following:*

13 “(n) *SPECIAL PROVISIONS FOR PRACTITIONERS CON-*  
14 *DUCTING CERTAIN RESEARCH WITH SCHEDULE I CON-*  
15 *TROLLED SUBSTANCES.—*

16 “(1) *IN GENERAL.—Notwithstanding subsection*  
17 *(f), a practitioner may conduct research described in*  
18 *paragraph (2) of this subsection with 1 or more*  
19 *schedule I substances in accordance with subparagraph*  
20 *(A) or (B) of paragraph (3) of this subsection.*

21 “(2) *RESEARCH SUBJECT TO EXPEDITED PROCES-*  
22 *DURES.—Research described in this paragraph is re-*  
23 *search that—*

24 “(A) *is with respect to a drug that is the*  
25 *subject of an investigational use exemption under*

1           *section 505(i) of the Federal Food, Drug, and*  
2           *Cosmetic Act; or*

3           “(B) is—

4               “(i) conducted by the Department of  
5           *Health and Human Services or the Depart-*  
6           *ment of Veterans Affairs; or*

7               “(ii) funded partly or entirely by a  
8           *grant, contract, cooperative agreement, or*  
9           *other transaction from the Department of*  
10          *Health and Human Services or the Depart-*  
11          *ment of Veterans Affairs.*

12          “(3) EXPEDITED PROCEDURES.—

13           “(A) RESEARCHER WITH A CURRENT  
14          *SCHEDULE I OR II RESEARCH REGISTRATION.—*

15               “(i) IN GENERAL.—*If a practitioner is*  
16           *registered to conduct research with a con-*  
17           *trolled substance in schedule I or II, the*  
18           *practitioner may conduct research under*  
19           *this subsection on and after the date that is*  
20           *30 days after the date on which the practi-*  
21           *tioner sends a notice to the Attorney Gen-*  
22           *eral containing the following information,*  
23           *with respect to each substance with which*  
24           *the practitioner will conduct the research:*

1                   “(I) The chemical name of the  
2 substance.

3                   “(II) The quantity of the sub-  
4 stance to be used in the research.

5                   “(III) Demonstration that the re-  
6 search is in the category described in  
7 paragraph (2), which demonstration  
8 may be satisfied—

9                   “(aa) in the case of a grant,  
10 contract, cooperative agreement,  
11 or other transaction, or intra-  
12 mural research project, by identi-  
13 fying the sponsoring agency and  
14 supplying the number of the  
15 grant, contract, cooperative agree-  
16 ment, other transaction, or  
17 project; or

18                   “(bb) in the case of an appli-  
19 cation under section 505(i) of the  
20 Federal Food, Drug, and Cosmetic  
21 Act, by supplying the application  
22 number and the sponsor of record  
23 on the application.

24                   “(IV) Demonstration that the re-  
25 searcher is authorized to conduct re-

1                   *search with respect to the substance  
2                   under the laws of the State in which  
3                   the research will take place.*

4                   “(ii) *VERIFICATION OF INFORMATION  
5                   BY HHS OR VA.*—*Upon request from the At-  
6                   torney General, the Secretary of Health and  
7                   Human Services or the Secretary of Vet-  
8                   erans Affairs, as appropriate, shall verify  
9                   information submitted by an applicant  
10                  under clause (i)(III).*

11                  “(B) *RESEARCHER WITHOUT A CURRENT  
12                  SCHEDULE I OR II RESEARCH REGISTRATION.*—

13                  “(i) *IN GENERAL.*—*If a practitioner is  
14                  not registered to conduct research with a  
15                  controlled substance in schedule I or II, the  
16                  practitioner may send a notice to the Attor-  
17                  ney General containing the information  
18                  listed in subparagraph (A)(i), with respect  
19                  to each substance with which the practi-  
20                  tioner will conduct the research.*

21                  “(ii) *ATTORNEY GENERAL ACTION.*—  
22                  *The Attorney General shall—*

23                  “(I) *treat notice received under  
24                  clause (i) as a sufficient application  
25                  for a research registration; and*

1                         “(II) not later than 45 days of re-  
2 ceiving such a notice that contains all  
3 information required under subparagraph  
4 (A)(i)—

5                         “(aa) register the applicant;  
6 or

7                         “(bb) serve an order to show  
8 cause upon the applicant in ac-  
9 cordance with section 304(c).

10                         “(4) ELECTRONIC SUBMISSIONS.—The Attorney  
11 General shall provide a means to permit a practi-  
12 tioner to submit a notification under paragraph (3)  
13 electronically.

14                         “(5) LIMITATION ON AMOUNTS.—A practitioner  
15 conducting research with a schedule I substance under  
16 this subsection may only possess the amounts of  
17 schedule I substance identified in—

18                         “(A) the notification to the Attorney Gen-  
19 eral under paragraph (3); or

20                         “(B) a supplemental notification that the  
21 practitioner may send if the practitioner needs  
22 additional amounts for the research, which sup-  
23 plemental notification shall include—

24                         “(i) the name of the practitioner;

1                   “(ii) the additional quantity needed of  
2                   the substance; and

3                   “(iii) an attestation that the research  
4                   to be conducted with the substance is con-  
5                   sistent with the scope of the research that  
6                   was the subject of the notification under  
7                   paragraph (3).

8                   “(6) IMPORTATION AND EXPORTATION REQUIRE-  
9                   MENTS NOT AFFECTED.—Nothing in this subsection  
10                  alters the requirements of part A of title III, regard-  
11                  ing the importation and exportation of controlled sub-  
12                  stances.”.

13                 (b) SEPARATE REGISTRATIONS NOT REQUIRED FOR  
14 ADDITIONAL RESEARCHER IN SAME INSTITUTION.—Sec-  
15 tion 302(c) of the Controlled Substances Act (21 U.S.C.  
16 822(c)) is amended by adding at the end the following:

17                 “(4) An agent or employee of a research institu-  
18                 tion that is conducting research with a controlled sub-  
19                 stance if—

20                 “(A) the agent or employee is acting within  
21                 the scope of the professional practice of the agent  
22                 or employee;

23                 “(B) another agent or employee of the insti-  
24                 tution is registered to conduct research with a  
25                 controlled substance in the same schedule;

1               “(C) the researcher who is so registered—

2                     “(i) informs the Attorney General of  
3                     the name, position title, and employing in-  
4                     stitution of the agent or employee who is  
5                     not separately registered;

6                     “(ii) authorizes that agent or employee  
7                     to perform research under the registration of  
8                     the registered researcher; and

9                     “(iii) affirms that any act taken by  
10                    that agent or employee involving a con-  
11                    trolled substance shall be attributable to the  
12                    registered researcher, as if the researcher  
13                    had directly committed the act, for purposes  
14                    of any proceeding under section 304(a) to  
15                    suspend or revoke the registration of the reg-  
16                    istered researcher; and

17                “(D) the Attorney General does not, within  
18                    30 days of receiving the information, authoriza-  
19                    tion, and affirmation described in subparagraph  
20                    (C), refuse, for a reason listed in section 304(a),  
21                    to allow the agent or employee to possess the sub-  
22                    stance without a separate registration.”.

23                (c) *SINGLE REGISTRATION FOR RELATED RESEARCH*  
24                SITES.—Section 302(e) of the Controlled Substances Act (21

1 U.S.C. 822(e)) is amended by adding at the end the fol-  
2 lowing:

3 “(4)(A) Notwithstanding paragraph (1), a person reg-  
4 istered to conduct research with a controlled substance  
5 under section 303(f) may conduct the research under a sin-  
6 gle registration if—

7 “(i) the research occurs exclusively on sites all of  
8 which are—

9 “(I) within the same city or county; and  
10 “(II) under the control of the same institu-  
11 tion, organization, or agency; and

12 “(ii) before commencing the research, the re-  
13 searcher notifies the Attorney General of each site  
14 where—

15 “(I) the research will be conducted; or

16 “(II) the controlled substance will be stored  
17 or administered.

18 “(B) A site described in subparagraph (A) shall be in-  
19 cluded in a registration described in that subparagraph  
20 only if the researcher has notified the Attorney General of  
21 the site—

22 “(i) in the application for the registration; or

23 “(ii) before the research is conducted, or before  
24 the controlled substance is stored or administered, at  
25 the site.

1       “(C) The Attorney General may, in consultation with  
2 the Secretary, issue regulations addressing, with respect to  
3 research sites described in subparagraph (A)—

4           “(i) the manner in which controlled substances  
5 may be delivered to the research sites;

6           “(ii) the storage and security of controlled sub-  
7 stances at the research sites;

8           “(iii) the maintenance of records for the research  
9 sites; and

10          “(iv) any other matters necessary to ensure effec-  
11 tive controls against diversion at the research sites.”.

12       (d) NEW INSPECTION NOT REQUIRED IN CERTAIN SIT-  
13 UATIONS.—Section 302(f) of the Controlled Substances Act  
14 (21 U.S.C. 822(f)) is amended—

15           (1) by striking “(f) The” and inserting “(f)(1)  
16 The”; and

17           (2) by adding at the end the following:

18           “(2)(A) If a person is registered to conduct research  
19 with a controlled substance and applies for a registration,  
20 or for a modification of a registration, to conduct research  
21 with a second controlled substance that is in the same sched-  
22 ule as the first controlled substance, or is in a schedule with  
23 a higher numerical designation than the schedule of the first  
24 controlled substance, a new inspection by the Attorney Gen-  
25 eral of the registered location is not required.

1       “(B) Nothing in subparagraph (A) shall prohibit the  
2 Attorney General from conducting an inspection that the  
3 Attorney General determines necessary to ensure that a reg-  
4 istrant maintains effective controls against diversion.”.

5       (e) *CONTINUATION OF RESEARCH ON SUBSTANCES*  
6 *NEWLY ADDED TO SCHEDULE I.*—Section 302 of the Con-  
7 trolled Substances Act (21 U.S.C. 822) is amended by add-  
8 ing at the end the following:

9       “(h) *CONTINUATION OF RESEARCH ON SUBSTANCES*  
10 *NEWLY ADDED TO SCHEDULE I.*—If a person is conducting  
11 research on a substance when the substance is added to  
12 schedule I, and the person is already registered to conduct  
13 research with a controlled substance in schedule I—

14           “(1) not later than 90 days after the scheduling  
15 of the newly scheduled substance, the person shall sub-  
16 mit a completed application for registration or modi-  
17 fication of existing registration, to conduct research  
18 on the substance, in accordance with regulations  
19 issued by the Attorney General for purposes of this  
20 paragraph;

21           “(2) the person may, notwithstanding sub-  
22 sections (a) and (b), continue to conduct the research  
23 on the substance until—

24           “(A) the person withdraws the application  
25 described in paragraph (1) of this subsection; or

1                 “(B) the Attorney General serves on the per-  
2                 son an order to show cause proposing the denial  
3                 of the application under section 304(c);

4                 “(3) if the Attorney General serves an order to  
5                 show cause as described in paragraph (2)(B) and the  
6                 person requests a hearing, the hearing shall be held on  
7                 an expedited basis and not later than 45 days after  
8                 the request is made, except that the hearing may be  
9                 held at a later time if so requested by the person; and

10                 “(4) if the person sends a copy of the application  
11                 described in paragraph (1) to a manufacturer or dis-  
12                 tributor of the substance, receipt of the copy by the  
13                 manufacturer or distributor shall constitute sufficient  
14                 evidence that the person is authorized to receive the  
15                 substance.”.

16                 (f) *TREATMENT OF CERTAIN MANUFACTURING ACTIVI-*  
17                 TIES AS COINCIDENT TO RESEARCH.—Section 302 of the  
18                 Controlled Substances Act (21 U.S.C. 822), as amended by  
19                 subsection (e), is amended by adding at the end the fol-  
20                 lowing:

21                 “(i) *TREATMENT OF CERTAIN MANUFACTURING AC-*  
22                 TIVITIES AS COINCIDENT TO RESEARCH.—

23                 “(1) IN GENERAL.—Except as provided in para-  
24                 graph (3), a person who is registered to perform re-  
25                 search on a controlled substance may perform manu-

1       *factoring activities with small quantities of that sub-*  
2       *stance, including activities described in paragraph*  
3       *(2), without being required to obtain a manufac-*  
4       *turing registration, if—*

5               “*(A) the activities are performed for the*  
6       *purpose of the research; and*

7               “*(B) the activities and the quantities of the*  
8       *substance involved in the activities are stated*  
9       *in—*

10              “*(i) a notification submitted to the At-*  
11       *torney General under section 303(l);*

12              “*(ii) a research protocol filed with an*  
13       *application for registration approval under*  
14       *section 303(f); or*

15              “*(iii) a notification to the Attorney*  
16       *General that includes—*

17              “*(I) the name of the registrant;*  
18       *and*

19              “*(II) an attestation that the re-*  
20       *search to be conducted with the small*  
21       *quantities of manufactured substance*  
22       *is consistent with the scope of the re-*  
23       *search that is the basis for the registra-*  
24       *tion.*

1           “(2) ACTIVITIES INCLUDED.—Activities per-  
2        mitted under paragraph (1) include—

3           “(A) processing the substance to create ex-  
4        tracts, tinctures, oils, solutions, derivatives, or  
5        other forms of the substance consistent with—

6           “(i) the information provided as part  
7        of a notification submitted to the Attorney  
8        General under section 303(l); or

9           “(ii) a research protocol filed with an  
10       application for registration approval under  
11       section 303(f); and

12           “(B) dosage form development studies per-  
13       formed for the purpose of requesting an inves-  
14       tigational new drug exemption under section  
15       505(i) of the Federal Food, Drug, and Cosmetic  
16       Act (21 U.S.C. 355(i)).

17           “(3) EXCEPTION REGARDING MARIHUANA.—The  
18       authority under paragraph (1) to manufacture sub-  
19       stances does not include the authority to grow mari-  
20       huana.”.

21           (g) TRANSPARENCY REGARDING SPECIAL PROCE-  
22       DURES.—Section 303 of the Controlled Substances Act (21  
23       U.S.C. 823), as amended by subsection (a), is amended by  
24       adding at the end the following:

1       “(o) TRANSPARENCY REGARDING SPECIAL PROCE-  
2 DURES.—

3           “(1) IN GENERAL.—If the Attorney General de-  
4 termines, with respect to a controlled substance, that  
5 an application by a practitioner to conduct research  
6 with the substance should be considered under a proc-  
7 ess, or subject to criteria, different from the process or  
8 criteria applicable to applications to conduct research  
9 with other controlled substances in the same schedule,  
10 the Attorney General shall make public, including by  
11 posting on the website of the Drug Enforcement Ad-  
12 ministration—

13           “(A) the identities of all substances for  
14 which such determinations have been made;

15           “(B) the process and criteria that shall be  
16 applied to applications to conduct research with  
17 those substances; and

18           “(C) how the process and criteria described  
19 in subparagraph (B) differ from the process and  
20 criteria applicable to applications to conduct re-  
21 search with other controlled substances in the  
22 same schedule.

23           “(2) TIMING OF POSTING.—The Attorney General  
24 shall make information described in paragraph (1)  
25 public upon making a determination described in

1       *that paragraph, regardless of whether a practitioner*  
2       *has submitted such an application at that time.”.*

3   **SEC. 4. RULEMAKING.**

4       (a) *INTERIM FINAL RULES.—The Attorney General—*  
5           (1) *shall, not later than 1 year of the date of en-*  
6           *actment of this Act, issue rules to implement this Act*  
7           *and the amendments made by this Act; and*  
8           (2) *may issue the rules under paragraph (1) as*  
9           *interim final rules.*

10     (b) *PROCEDURE FOR FINAL RULE.—*

11       (1) *EFFECTIVENESS OF INTERIM FINAL RULES.—*  
12       *A rule issued by the Attorney General as an interim*  
13       *final rule under subsection (a) shall become imme-*  
14       *diately effective as an interim final rule without re-*  
15       *quiring the Attorney General to demonstrate good*  
16       *cause therefor, notwithstanding subparagraph (B) of*  
17       *section 553(b) of title 5, United States Code.*

18       (2) *OPPORTUNITY FOR COMMENT AND HEAR-*  
19       *ING.—An interim final rule issued under subsection*  
20       *(a) shall give interested persons the opportunity to*  
21       *comment and to request a hearing.*

22       (3) *FINAL RULE.—After the conclusion of such*  
23       *proceedings, the Attorney General shall issue a final*  
24       *rule to implement this Act and the amendments made*

1       *by this Act in accordance with section 553 of title 5,*  
2       *United States Code.*

3   **SEC. 5. PENALTIES.**

4       (a) *IN GENERAL.*—Section 401(b)(1) of the Controlled  
5   *Substances Act (21 U.S.C. 841(b)(1)) is amended—*

6               (1) *in subparagraph (A)(vi), by inserting “or a*  
7       *fentanyl-related substance” after “any analogue of N-*  
8       *phenyl-N-[1-(2-phenylethyl)-4-piperidinyl]*  
9       *propanamide”; and*

10              (2) *in subparagraph (B)(vi), by inserting “or a*  
11       *fentanyl-related substance” after “any analogue of N-*  
12       *phenyl-N-[1-(2-phenylethyl)-4-piperidinyl]*  
13       *propanamide”.*

14       (b) *IMPORTATION AND EXPORTATION.*—Section  
15   *1010(b) of the Controlled Substances Import and Export*  
16   *Act (21 U.S.C. 960(b)) is amended—*

17              (1) *in paragraph (1)(F), by inserting “or a*  
18       *fentanyl-related substance” after “any analogue of N-*  
19       *phenyl-N-[1-(2-phenylethyl)-4-piperidinyl]*  
20       *propanamide”; and*

21              (2) *in paragraph (2)(F), by inserting “or a*  
22       *fentanyl-related substance” after “any analogue of N-*  
23       *phenyl-N-[1-(2-phenylethyl)-4-piperidinyl]*  
24       *propanamide”.*

1   **SEC. 6. APPLICABILITY; OTHER MATTERS.**

2       (a) *IN GENERAL.*—Irrespective of the date on which  
3   the rules required by section 4 are finalized, the amend-  
4   ments made by this Act apply beginning as of the enact-  
5   ment of this Act.

6       (b) *RULE OF CONSTRUCTION.*—Nothing in the amend-  
7   ments made by this Act may be construed as evidence that,  
8   in applying sections 401(b)(1) and 1010(b) of the Con-  
9   trolled Substances Act (21 U.S.C. 841(b)(1) and 960(b))  
10   with respect to conduct occurring before the date of the en-  
11   actment of this Act, a fentanyl-related substance (as defined  
12   by such amendments) is not an analogue of N-phenyl-N-  
13   [1-(2-phenylethyl)-4-piperidinyl] propanamide.

14       (c) *SENSE OF CONGRESS.*—The Congress agrees with  
15   the interpretation of the Controlled Substances Act (21  
16   U.S.C. 801 et seq.) in *United States v. McCray*, 346 F.  
17   Supp. 3d 363 (2018).

**Union Calendar No. 47**

118TH CONGRESS  
1ST SESSION

**H. R. 467**

**[Report No. 118-67, Part I]**

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**A BILL**

To amend the Controlled Substances Act with respect to the scheduling of fentanyl-related substances, and for other purposes.

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MAY 17, 2023

Reported from the Committee on Energy and Commerce  
with an amendment

MAY 17, 2023

Committee on the Judiciary discharged; committed to the  
Committee of the Whole House on the State of the  
Union and ordered to be printed